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1    **Estimation of bladder contractility from intravesical pressure-volume measurements.**

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21

22    Short title. Measurement of bladder contractility

23

## 24 **Abstract**

25 *Aim.* To describe parameters from urodynamic pressure recordings that describe urinary  
26 bladder contractility through the use of principles of muscle mechanics.

27 *Methods.* Subtracted detrusor pressure and voided flow were recorded from patients  
28 undergoing filling cystometry. The isovolumetric increase of detrusor pressure,  $p$ , of a  
29 voluntary bladder contraction before voiding was used to generate a plot of  $(dp/dt)/p$  vs  $p$ .  
30 Extrapolation of the plot to the  $y$ -axis and the  $x$ -axis generated a contractility parameter,  $v_{CE}$   
31 (the maximum rate of pressure development) and the maximum isovolumetric pressure,  $p_0$ ,  
32 respectively. Similar curves were obtained in *ex vivo* pig bladders with different  
33 concentrations of the inotropic agent carbachol and shown in a supplement.

34 *Results.* Values of  $v_{CE}$ , but not  $p_0$ , diminished with age in female subjects.  $v_{CE}$  was most  
35 significantly associated with the 20-80% duration of isovolumetric contraction  $t_{20-80}$ ; and a  
36 weaker association with maximum flow rate and BCI in women.  $p_0$  was not associated with  
37 any urodynamic variable in women, but in men was with  $t_{20-80}$  and isovolumetric pressure  
38 indices.

39 *Conclusions.* The rate of isovolumetric subtracted detrusor pressure ( $t_{20-80}$ ) increase shows a  
40 very significant association with indices of bladder contractility as derived from a derived  
41 force-velocity curve. We propose that  $t_{20-80}$  is a detrusor contractility parameter (DCP).

42

## 43 Introduction

44 To void completely the bladder must generate sufficient intravesical pressure to overcome  
45 outflow tract resistance and sustain this to allow complete voiding. However, the bladder  
46 may generate excessively large pressures that can impact on upper tract integrity or may  
47 exhibit an underactive phenotype, resulting in incomplete bladder emptying.<sup>1</sup> There is a  
48 clinical need to identify the causes of abnormal voiding to allow proper management of these  
49 conditions and identify people at risk if being considered for surgery to improve voiding. In  
50 particular, men who require relief from voiding lower urinary tract symptoms (LUTS) may  
51 not benefit from surgery if impaired bladder contractility is the cause. Likewise, women may  
52 be at risk of voiding difficulty after surgery to treat stress urinary incontinence, even if there  
53 were no prior voiding LUTS.

54

55 A rise of detrusor pressure results from increased bladder wall tension generated by  
56 contracting detrusor smooth muscle. However, several lower urinary tract properties  
57 contribute to voiding: the magnitude of urethral resistance; the strength and duration of  
58 detrusor smooth muscle contraction; and initial bladder volume which by Laplace's Law is an  
59 inverse function of detrusor pressure. Of fundamental interest is whether or not impaired or  
60 enhanced detrusor muscle contractile function contributes to abnormal voiding. Several  
61 urodynamic-based definitions of bladder contractility have been defined, e.g. a bladder  
62 contractility index (BCI)<sup>2</sup>, Watts factor<sup>3</sup> and bladder outlet relation,<sup>4</sup> but these are also  
63 influenced by outflow tract properties. Furthermore, such parameters are poorly validated in  
64 women. Therefore, it is desirable to define a gender-neutral urodynamic-based parameter  
65 that reflects physiological bladder contractility.

66

67 Smooth and cardiac muscle exhibit a property that contractile strength depends not only on  
68 resting fibre length but also on an intrinsic inotropic state. The latter is an alteration of  
69 contractile performance independent of resting length and physiologically is defined as a  
70 change of contractility: this term therefore has a specific meaning for isometric (muscle  
71 tension developed at constant length) or isotonic (muscle shortening at constant load)  
72 contractions. Changes of contractility have been demonstrated in the intact heart and isolated  
73 cardiac preparations<sup>5-7</sup> and evaluated in the bladder.<sup>8,9</sup> To identify patients with reduced  
74 cardiac contractility (heart failure), it was important to identify parameters independent of  
75 preload and afterload, as these affect muscle fibre length and contractile strength.<sup>5</sup> The  
76 usefulness of indices derived from the rise of isovolumetric ventricular pressure ( $P$ ) in the  
77 cardiac cycle to define cardiac contractility is validated, especially from a plot of  $(dP/dt).P^{-1}$  as  
78 a function of  $P$  itself.<sup>10,11</sup> These indices are the extrapolated fits of this relation to: a) the  $y$ -axis  
79  $(dP/dt).P^{-1}$ -axis) which represents maximum velocity of muscle shortening,  $v_{CE}$ ; b) the  $x$ -axis  
80 ( $P$ -axis) which represents maximum isovolumetric pressure,  $P_0$ . The indices reflect the action  
81 of positive and negative inotropic agents and thus changes to cardiac contractility. However,  
82 this analysis has not been applied to smooth muscle-lined organs. We used urodynamic  
83 recordings of bladder contractile function to obtain a practical measure of true bladder  
84 contractility based on the above cardiac principles and how this may be used to understand  
85 different disorders of bladder function.

86

## 87 Materials and Methods

88 *Data collection.* Anonymised data were obtained from 49 patients attending the urodynamic  
89 clinic at Southmead Hospital, North Bristol NHS Trust. Inclusion criteria were patients listed  
90 for routine or video urodynamics and with voided volumes greater than 100 ml. Pressure  
91 readings on both abdominal and intravesical lines were obtained following ICS guidelines.<sup>12</sup>  
92 Exclusion criteria were: evidence of a bladder diverticulum which would prevent a constant-  
93 volume contraction (four); voiding achieved mainly with additional abdominal straining  
94 (five), leaving 40 records (21 male, 19 female patients) for analysis. Data were anonymised  
95 and collected according to standard departmental protocols as part of urodynamic testing.  
96 Ethical approval was obtained from the regional research ethics committee.

97

98 *Urodynamic measurements in subjects.* Subjects emptied their bladder before urodynamics.  
99 An Aquarius urodynamics system (Laborie; Mississauga, Canada) used water-filled catheters  
100 to record infused volume as a function of time and values of abdominal and intravesical  
101 pressures to obtain calculated detrusor pressure,  $P_{\text{det}}$ , in accordance with ICS Good  
102 Urodynamics Practices.<sup>12</sup> Flow during voiding was measured directly. Analogue  
103 measurements were digitised at 50 kHz and stored on the urodynamics system computer.  
104 For all other aspects, the manufacturer's default values were used for data acquisition, namely  
105 0.8 seconds delay time added to the pressure trace to align it with the flow-meter recording.  
106 Data were retrieved from these files at 0.05 kHz for subsequent analysis. Figure 1A, B shows  
107 the different measurements made from urodynamic recordings. Derived indices were: the  
108 maximum rate of  $P_{\text{det}}$  change during isovolumetric contraction,  $dP_{\text{det}}/dt_{\text{max}}$ ; isovolumetric  
109 contraction duration from 20% to 80%  $\Delta P_{\text{isv}}$  ( $t_{80-20}$ ); bladder contractility index (BCI;  
110  $P_{\text{det}}Q_{\text{max}}+5Q_{\text{max}}$ ) and bladder outflow obstruction index (BOOI;  $P_{\text{det}}Q_{\text{max}}-2Q_{\text{max}}$ ).

111

112 *Smoothing pressure traces.*  $P_{\text{det}}$  measurements were made during isovolumetric bladder  
 113 contractions, between the onset of a rise of detrusor pressure and the beginning of flow,  $Q$   
 114 (Figure 2A). The base-line (pre-contraction) value of  $P_{\text{det}}$  was subtracted from values during  
 115 contraction, henceforth called  $P$ . The dependent variable in the final analysis,  $(dP/dt)/P^{-1}$  –  
 116 see supplementary material - is significantly influenced by artifactual pressure fluctuations  
 117 and  $P$ -data were smoothed by calculating running averages ( $RA$ ), equation 1, with  $n=10, 20,$   
 118  $50$  or  $100$ , i.e. averaging occurred over  $0.2, 0.4, 1.0$  or  $2.0$  s:

$$119 \quad RA_n = (x_1 + x_2 + \dots + x_n)/n, \quad RA_{n+1} = (x_2 + x_3 + \dots + x_{n+1})/n, \quad RA_{n+2} = (x_3 + x_4 + \dots + x_{n+2})/n, \dots \quad 1$$

120 In this analysis a value of  $n=20$  was used, i.e. a value of  $0.4$  s running average time was  
 121 chosen that imposed only a 2% slowing of the pressure trace.

122

123 *Data analysis.* The supplementary data describe the rationale for estimating a contractility  
 124 parameter of an isovolumetrically-contracting spheroid organ. A plot of  $(dP/dt).P^{-1}$  vs  $P$  yields  
 125 a phase loop of which the declining portion at higher  $P$ -values is described by a hyperbolic  
 126 function that intercepts the  $y$ -axis (i.e. at  $P=0$ ) to yield a value for the maximum velocity of  
 127 contractile element shortening ( $v_{\text{CE}}$ ). An increase of  $v_{\text{CE}}$  is interpreted as enhanced bladder  
 128 contractility. The intercept on the  $x$ -axis is the maximal isovolumetric pressure,  $P_0$ , that would  
 129 have been achieved had the outflow tract not opened. A value of bladder wall tension,  $T_0$ , was  
 130 calculated from the Laplace equation:  $T_0 = P_0.r/2$ ; where  $r$  is bladder radius, calculated from  
 131 the filling volume,  $V$ , at initiation of voiding: i.e.  $r = \sqrt[3]{\frac{0.75V}{\pi}}$ . Values of  $v_{\text{CE}}$  and  $P_0$  were  
 132 compared to indices as measured or calculated from urodynamic traces (Figure 1) to  
 133 determine where the best associations were observed. For isolated pig bladders data values  
 134 of  $v_{\text{CE}}$  and  $P_0$  were plotted as a function of the carbachol concentration.

135

136 *Data presentation and statistical analyses.* Group values are represented as medians (25, 75%  
137 interquartiles). Associations between variables were calculated from a Spearman's  
138 correlation coefficient,  $\rho$ , whose significance was tested by calculation of a  $t$ -value from  $\rho$   
139 using  $t = \rho \sqrt{\frac{n-2}{1-\rho^2}}$ , with subsequent estimation of  $p$ -values. Differences between sets were  
140 tested by ANOVA, with *post hoc* Bonferroni multiple comparisons. In total, 10 (see Results)  
141 separate hypotheses were tested for a significant association between  $v_{CE}$  or  $P_0$  and a  
142 particular urodynamic variable. To minimise the chance of a spurious level of significance  
143 arising from these multiple tests the  $p$ -value for significance was reduced from one of  $p < 0.05$   
144 by a correction to  $p < 0.005$  (i.e.  $p < 0.05/10$ ). A least-squares iterative fitting program  
145 (KaleidaGraph, Synergy Inc) was used to fit linear and non-linear functions to data sets.  
146



## 147 Results

148 *Patient urodynamic characteristics and contractility indices.* Table 1 shows urodynamic-  
149 derived values of lower urinary tract function (figure 1). All variables were similar in males  
150 and females, except: in females calculated BOOI was smaller. Values of the contractility indices  
151  $v_{CE}$  and  $P_0$ , as well as calculated  $T_0$ , are also shown in table 1 and were also similar in males  
152 and females.

153

154 *Smoothing of pressure waveforms.* Figure 2A shows an example of the rise of subtracted  
155 detrusor pressure in the isovolumetric phase for an unsmoothed ( $P$ ) trace and smoothed  
156  $n=10, 20, 50$  or  $100$  (eq 1, Methods) i.e. averaging over 0.2, 0.4, 1.0 or 2.0 s for data recorded  
157 at 0.05 kHz. Traces are shown as  $P, P_{10}, P_{20}, P_{50}$  and  $P_{100}$ . The extent of time delay is shown  
158 in Figure 2B where values of  $P$  are plotted against  $P_n$ . For no introduced time-delay the slope  
159 of the line would be unity. From 28 sample traces the slopes were 1.007 [0.999, 1.104],  
160 1.020 [1.000, 1.033], 1.044 [1.007, 1.083] for averaging over 0.2 ( $n=10$ ), 0.4 ( $n=20$ ), 1.0  
161 ( $n=50$ ) and 2.0 ( $n=100$ ) s, respectively. Heuristically a value of  $n=20$  was chosen for onward  
162 analysis, which would introduce an error of 2% [0.0 – 3.3%] in traces and consequent  
163 derived values of  $v_{CE}$ .

164

165 *Relation of  $v_{CE}$  to urodynamic variables.*  $v_{CE}$  values were negatively associated with age for  
166 females ( $p=0.03$ ;  $n=19$ ; 22-85 years) but not for males ( $n=21$ ; 22-84 years) subjects. For  
167 urodynamic variables there were significant negative associations for female and male  
168 subjects with  $t_{20-80}$  ( $p=0.000002$  and  $0.002$ , respectively;  $r=-0.636$  and  $-0.861$ ). Positive  
169 associations for females only were also shown for  $Q_{max}$  and BCI (both  $p=0.002$ ;  $r=0.668$  and  
170  $0.663$ , respectively). There were no significant associations for other urodynamic variables,  
171 i.e.  $\Delta P_{det}$ ,  $P_{det}Q_{init}$ ,  $\Delta P_{isv}$ ,  $P_{det}Q_{max}$ ,  $P_{det}$  time or  $Q$  time, nor the calculated index BOOI for either

172 gender. Figure 3A-D shows the relationships between  $v_{CE}$  and age,  $t_{20-80}$ ,  $Q_{max}$  or BCI for  
173 female and male subjects.

174

175 *Relation of  $P_0$  to urodynamic variables.* There was no association of  $P_0$  with age for subjects of  
176 either gender. A different pattern of associations was observed for  $P_0$  with urodynamic  
177 variables compared to  $v_{CE}$ . For female subjects no urodynamic variable showed a significant  
178 relationship with  $P_0$ . For male subjects  $P_0$  was positively associated with  $\Delta P_{det}$ ,  $P_{det}Q_{init}$ ,  $\Delta P_{isv}$   
179 and  $P_{det}Q_{max}$  (all  $p<0.00001$ ;  $r=0.893, 0.839, 0.917, 0.816$  respectively).  $P_0$  was also negatively  
180 associated with  $t_{20-80}$  ( $p=0.005$ ;  $r=-0.694$ ) and BOOI ( $p<0.0001$ ;  $r=-0.751$ ), but not for  $P_{det}$  time,  
181  $Q_{max}$ ,  $Q$  time nor BCI. Figure 3E shows the association between  $P_0$  and  $\Delta P_{isv}$ .

182

## 183 Discussion

184 *Description of bladder contractility.* Transformation of the isovolumetric contraction phase  
185 allows calculation of bladder wall contractility from pressure urodynamic traces in patients.  
186 Key to the analysis is estimation of muscle contractility from the intercept of a force-velocity  
187 curve with the  $y$ -axis,  $v_{CE}$ , which represents the maximum, unloaded velocity of shortening.  
188 Developed for tension measurements in skeletal muscle,<sup>12</sup> the analysis of  $v_{CE}$  was then applied  
189 to cardiac and smooth muscle preparations.<sup>13,14</sup> Its verification in isolated smooth muscle  
190 cells<sup>9,15</sup> further demonstrated that the relationship is a general feature of muscle cells.  $v_{CE}$  is  
191 related to crossbridge turnover and myosin ATPase muscle activity, and positive inotropic  
192 interventions increase the value.<sup>14</sup> The principle of generating force-velocity curves was  
193 extrapolated to isovolumetric hollow organs such as the heart, where pressure is proportional  
194 to tangential wall tension, and it was shown that addition of myocardial inotropic agents  
195 altered  $v_{CE}$  values in a consistent manner.<sup>11</sup> The method therefore generates a contractility  
196 index that derives from changes to the speed of contraction of the bladder wall.

197  
198 We have shown that force-velocity curves may also be generated from isovolumetric phases  
199 of voiding bladder contractions, and validated the approach in pig bladders (see supplement),  
200 showing that an inotropic agent, carbachol, altered  $v_{CE}$ . Moreover, comparison of  $v_{CE}$  values  
201 with urodynamic variables showed that the strongest association was with time-dependent  
202 variables of the isovolumetric rise of pressure, before voiding begins, i.e. the time from 20-  
203 80% ( $t_{20-80}$ ) of this phase. We propose this interval is a superior measure of bladder  
204 contractility as derived from urodynamic traces. Moreover, this association was independent  
205 of gender, unlike the case for urodynamic variables such as BCI, which is verified only for  
206 men. A parallel increase of maximum isovolumetric pressure,  $P_0$ , and contractility would be  
207 anticipated in an isovolumetric system but not in a voiding bladder, as in the latter the energy

208 of contraction would also be dissipated in causing urine flow at the expense of a continuous  
209 rise of pressure. Data from an isolated, arterially perfused pig bladder validated the approach  
210 by showing that addition of a contractile agonist, carbachol in rising concentrations, increased  
211  $v_{CE}$ , as well as both the maximum rate of increase of pressure and maximum pressure itself.

212

213 Although there was a significant association between  $v_{CE}$  and  $P_0$  in the human bladder these  
214 two derived variables measure different aspects of bladder function. Values of  $v_{CE}$ , reflecting  
215 different states of contractility and with units of reciprocal time, were strongly associated  
216 with the time course of the increase of isovolumetric pressure, as may be expected, but not  
217 with any urodynamic pressure parameter.  $P_0$  is a measure of steady-state tension and is  
218 determined also by the number of muscle fibres recruited to enable a detrusor contraction, as  
219 well as the contractility of individual muscle fibres.  $P_0$  showed stronger associations with  
220 pressure urodynamic indices than with time-dependent indices of isovolumetric pressure and  
221 is a less reliable estimation of detrusor muscle contractility itself.

222

223 *Gender differences.* There were no gender differences in  $v_{CE}$  or  $P_0$ , nor in individual  
224 urodynamic variables, except BOOI: table 1. However, there were striking differences  
225 between associations of  $v_{CE}$  or  $P_0$  with urodynamic variables. For  $v_{CE}$  values, an age-  
226 dependent decline was present in females but not males, suggesting a decline of detrusor  
227 contractility with age in women. There was also a significant association of  $v_{CE}$  with  $Q_{max}$ , and  
228 its derived variable BCI, in females but not males: thus in males factors other than muscle  
229 contractility, e.g. outflow tract resistance, influence the rate of urine flow.

230

231 *Limitations to the generation of force-velocity curves in an isovolumetric bladder.* i) Pressure in  
232 a hollow organ is proportional to tangential wall tension if it is truly isovolumetric and wall

233 thickness is unchanging during contraction.<sup>16</sup> Thus, patients with evident bladder diverticula  
234 should be excluded from the analysis, as chamber volume would change during these  
235 otherwise isovolumetric contractions. ii) In principle, contractility estimates are based on  
236 changes to muscle or wall tension. In different patients, bladder volume during isovolumetric  
237 contraction was not very variable and the relationship between  $\Delta P_{\text{isv}}$  and calculated wall  
238 tension was very significant ( $\rho=0.987$ ,  $n=40$ ,  $p<0.000001$ ). Thus, changes of isovolumetric  
239 pressure during voiding are proportional to those of wall tension. iii) The absolute value of  
240 the force-velocity curve intercept with the  $y$ -axis to estimate  $v_{\text{CE}}$  also depends on wall  
241 stiffness,  $k$ , and it was assumed to be unchanging during the contraction. iv) The urodynamic  
242 system that was used imposes an in-built delay of 0.8 s between the pressure and flow values,  
243 so that this final period of the isovolumetric pressure curve would be unreliable for analysis  
244 and was therefore not used. v) All analyses were performed on voluntary contractions - it will  
245 be of further interest to also analyse involuntary detrusor contractions.

246

247 Conclusions. The maximum velocity of an unloaded detrusor contraction,  $v_{\text{CE}}$ , shows a very  
248 close association with the urodynamic 20-80% time of the isovolumetric pressure  
249 contraction,  $t_{20-80}$ , we propose calling the detrusor contractility parameter (DCM).  $v_{\text{CE}}$  is an  
250 index of muscle contractility as verified in parallel pig bladders experiments. This method  
251 offers considerable advantages over current urodynamic estimates of contractility such as  
252 BCI, which is a flow- and volume-dependent variable and verified only for men. Its  
253 measurement can contribute to the debate regarding the concept of bladder underactivity as a  
254 true decline of bladder contractile function, rather than a degradation of other changes to the  
255 lower urinary tract that can impact on the magnitude and extent of voiding function.

256

257 List of abbreviations (see also legend to figure 1 for derived parameters and variables)

258  $P_{\text{det}}$  subtracted detrusor pressure

259  $P$   $P_{\text{det}} - (\text{baseline } P_{\text{det}} \text{ prior to isovolumetric pressure rise}).$

260  $P_0$  calculated maximum isovolumetric pressure

261  $v_{\text{CE}}$  calculated maximum velocity of contractile element shortening

262 BCI bladder contractility index

263 DCP detrusor contractility parameter

264  $T$  bladder wall tension

265  $Q$  flow

266 BOOI bladder outflow obstruction index

267  $t_{20-80\%}$  duration of 20-80% isovolumetric contraction interval

268  $k$  bladder wall stiffness

269

270 Notes: upper case  $P$  denotes pressure, rather than lower case  $p$  as often used in urodynamics

271 recordings. This is to avoid confusion with the p-value used to denote levels of statistical

272 significance. Lower case  $v$  is used to denote velocity to avoid confusion with  $V$  as often used in

273 urodynamics for volume.

274

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318



319 Figure 1. Urodynamic recordings of subtracted detrusor pressure,  $P_{\text{det}}$ , and flow,  $Q$ , from  
 320 which variables were measured. A: recordings over the time-frame of an entire bladder  
 321 contraction defining values of  $\Delta P_{\text{det}}$ ,  $P_{\text{det max}}$ ,  $P_{\text{det time}}$  and  $Flow time$ . The baseline  $P_{\text{det}}$  value is  
 322 also shown. B: faster time scale of the rise of the  $P_{\text{det}}$ -transient defining values of  $\Delta P_{\text{isv}}$ ,  
 323  $P_{\text{det}Q_{\text{init}}}$ ,  $P_{\text{det}Q_{\text{max}}}$  and  $Q_{\text{max}}$ . Also shown are the pressure values for 20 and 80% of  $\Delta P_{\text{isv}}$ , from  
 324 which the 20-80%  $P_{\text{isv}}$  time was calculated.

325

326 Figure 2. Smoothing protocols of pressure rises. A: The rising phase of a voiding  $P_{\text{det}}$ -  
 327 transient, either unsmoothed ( $P$ ) or smoothed with greater running average intervals (See  
 328 Methods, equation 1). The particular record was sampled at 50 Hz so that running averages  
 329 over 10, 20, 50 or 100 points generated traces averaged over 0.2, 0.4, 1.0 or 2.0 s ( $P_n = P_{10}$ -  
 330  $P_{100}$ ). B: Plots of running averaged traces ( $P_n$ ) as a function of the unaveraged trace ( $P$ ). The  
 331 line of identity is also drawn.

332

333 Figure 3. Derived contractility measures vs urodynamic variables. Plots of  $v_{\text{CE}}$  as a function of  
 334 age (part A);  $t_{20-80\%}$  (part B);  $Q_{\text{max}}$  (part C) and BCI (part D). The relation between  $P_0$  and  $\Delta P_{\text{isv}}$   
 335 is shown in part E. Data are shown separately for male participants (closed squares) and  
 336 female participants (open squares).

337

338 Table I. Demographic and urodynamic data, and derived contractility indices, of subjects  
 339 contributing to the study. Median data with 25, 75% interquartiles,  $n$ =number of patients

Figure 1

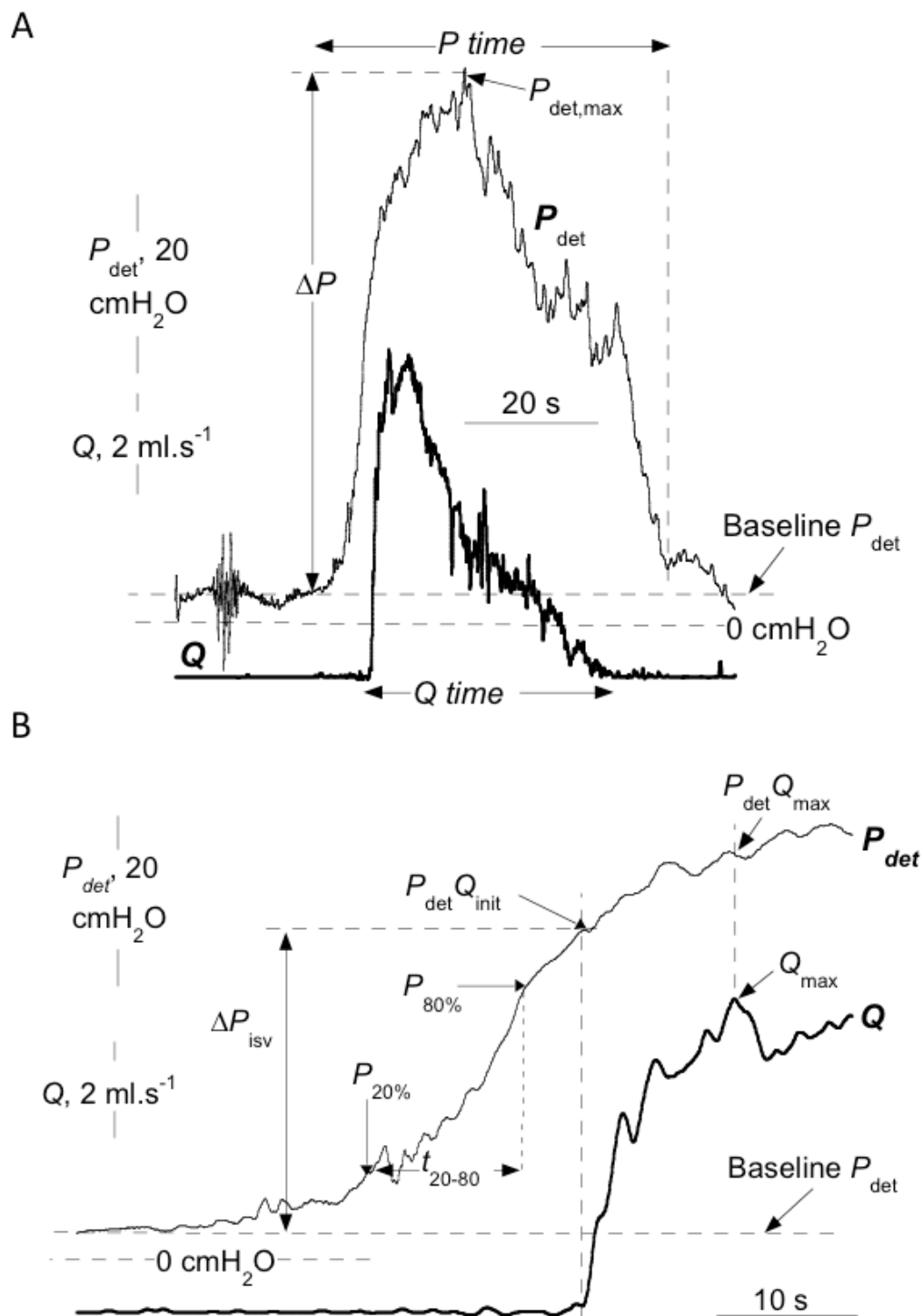
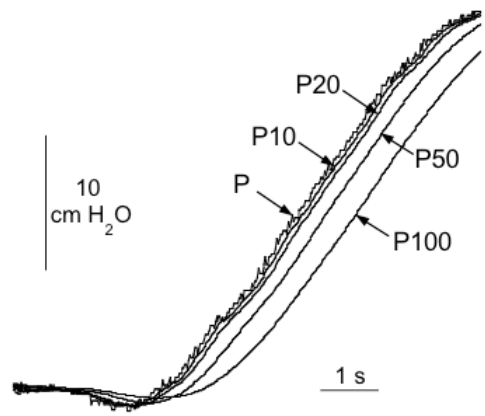


Figure 2

A



B

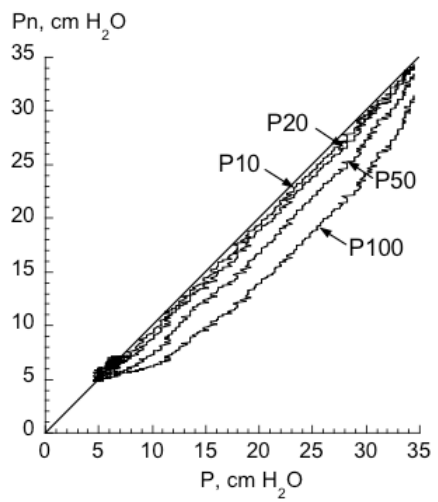


Figure 3

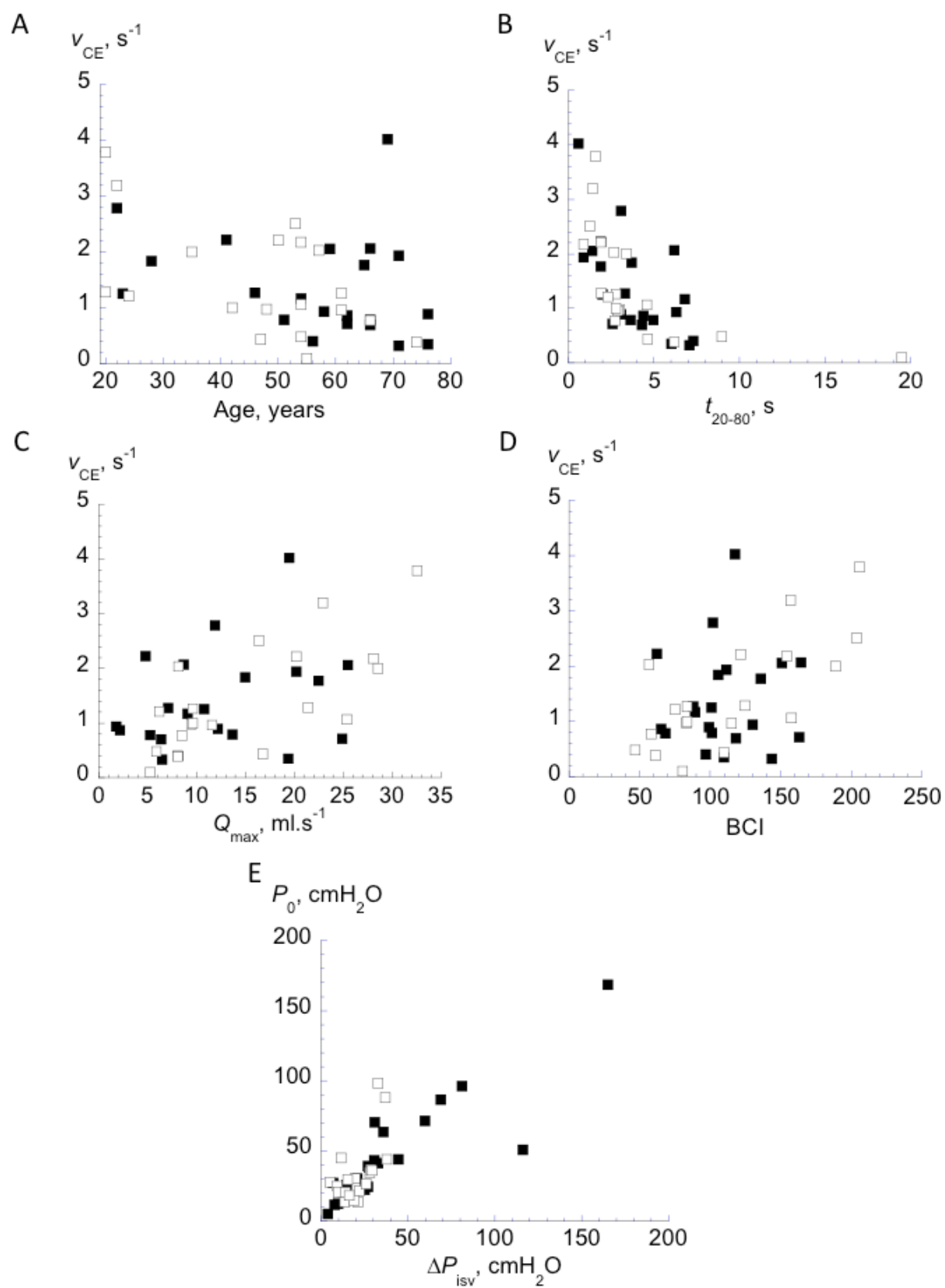


Table 1. Demographic and urodynamic data, and derived contractility indices, of subjects contributing to the study. Median data with 25, 75% interquartiles,  $n$ =number of patients.

	All data ( $n=40$ )	Male ( $n=21$ )	Female ( $n=19$ )
Clinical characteristics			
Age, years	54.5 [45, 65]	62 [51, 66]	53 [39, 56]
$\Delta p$ , cm H <sub>2</sub> O	38.5 [20.8, 55.6]	39.3 [24.7, 60.3]	34.5 [18.0, 44.8]
$p @ Q_{init}$ , cm H <sub>2</sub> O	33.9 [19.7, 41.1]	34.0 [22.7, 56.7]	31.0 [17.1, 36.8]
$\Delta p_{isovol}$ , cm H <sub>2</sub> O	25.4 [13.8, 32.8]	27.5 [14.9, 44.5]	19.4 [13.7, 28.4]
$Q_{max}$ , ml.s <sup>-1</sup>	11.2 [7.9, 20.2]	10.8 [6.5, 19.4]	11.6 [8.4, 22.2]
$p @ Q_{max}$ , cm H <sub>2</sub> O	38.1 [22.7, 48.3]	41.9 [30.7, 54.4]	35.6 [19.2, 43.6]
$BCI$	108 [84, 138]	106 [97, 130]	110 [78, 156]
$BOOI$	9.6 [-13.2, 32.3]	20.0 [-11.1, 40.1]	-0.8 [-15.5, 16.7] *
$V @ Q_{init}$ , ml	250 [225, 282]	229 [203, 249]	282 [250, 313]
$p$ duration, s	53.0 [25.9, 76.6]	52.0 [26.2, 90.0]	53.0 [29.1, 72.5]
$Q$ duration, s	31.8 [21.7, 42.0]	28.0 [23.0, 37.0]	34.0 [19.5, 51.5]
$t_{0-100\%}$ , s	8.5 [5.2, 12.0]	8.7 [4.5, 12.3]	8.1 [5.4, 11.7]
$t_{10-90\%}$ , s	4.0 [2.9, 6.7]	5.2 [3.3, 7.6]	3.5 [2.7, 5.4]
$t_{20-80\%}$ , s	3.0 [1.9, 4.7]	3.6 [2.0, 6.0]	2.8 [1.9, 4.0]
$dP/dt_{max, isovol}$ , cm H <sub>2</sub> O.s <sup>-1</sup>	6.8 [4.8, 9.8]	7.2 [4.7, 9.1]	6.8 [4.3, 10.3]
Contractility indices, voiding contractions			
$V_{CM}$ , s <sup>-1</sup>	1.19 [0.78, 2.04]	1.17 [0.78, 1.94]	1.21 [0.87, 2.11]
$p_0$ , cm H <sub>2</sub> O	30.2 [21.5, 44.3]	39.1 [24.6, 63.5]	27.9 [20.6, 36.4]
$T_0$ , N	68.9 [44.1, 89.1]	69.2 [52.2, 110.2]	59.3 [38.7, 74.3]

348 On-line Supplement.

349 Rationale of phase-loop analysis to calculate  $v_{CE}$  and  $P_0$ .

350 The use of  $(dP/dt).P^{-1}$  at zero load as an index of contractility follows from the Hill model for  
351 muscle contraction. A contractile, or force-generating, element (CE) lies in series with a series  
352 elastic component (SEC); the whole lying in parallel to a parallel elastic component (PEC).  
353 During an isometric contraction the rate,  $v$ , of CE shortening is equal to the rate of SEC  
354 lengthening so that:  $v_{CE} = v_{SEC} = |dl_{CE}/dt| = |dl_{SEC}/dt|$ : where  $l$  is the length of CE or SEC. The  
355 term  $v_{SEC}$  represents the rate of change of wall stress,  $s$ , per unit amount of stress, and is  
356 normalised to a stiffness constant for the tissue,  $k$ . Thus,  $v_{CE} = v_{SEC} = \frac{ds}{dt} \cdot (k.s)^{-1}$ . Laplace's  
357 Law shows that intravesical pressure,  $P$ , is proportional to wall stress under isovolumetric  
358 conditions, with an arbitrary constant,  $X$ , so that:  $v_{CE} = \left( X \frac{dP}{dt} \right) \cdot (k.X.P)^{-1} = \frac{dP}{dt} \cdot (k.P)^{-1}$ . A plot of  
359  $(dP/dt).P^{-1}$  as a continuous function of  $P$  throughout an isovolumetric contraction will yield a  
360 force (pressure)-velocity plot that will yield an extrapolated maximum value of  $v_{CE}$  at zero  
361 load (where the plot intercepts the  $(dP/dt).P^{-1}$  axis at  $P=0$ ) and an extrapolated maximum  
362 value of  $P$  ( $P_0$ ), where the plot intercepts the  $P$  axis as  $v_{CE}$  is zero.

363

364 Several assumptions and calculations were included.

365 1. It is assumed that the bladder is ellipsoid or spherical during isometric contraction and the  
366 rate of intravesical pressure change by Laplace's Law is proportional to the rate of contractile  
367 element shortening  
368 2. Contractile component tension is equal to series elastic (SEC) tension soon after activation.  
369 However, appreciation of PEC stress-strain properties is also needed as the PEC carries part of  
370 the resting tension and is only transferred to the SEC/CC fraction when CC shortening has  
371 been carried out. This may be accounted for by subtracting the absolute value of resting

372 tension (pressure) from values during isometric contraction and this was done in the  
373 measurement presented here.<sup>1</sup>

374 3. The series elastic tension is scaled by a stiffness parameter,  $k$ , that is assumed to be  
375 constant throughout. Whether  $k$  is equivalent or proportional to the reciprocal of bladder  
376 compliance during filling has not been determined and has been added to the 'limitations'  
377 section. However, this limitation will apply to any measurement of intravesical pressure  
378 during a contraction.

379

380 Supplementary Figure 1 shows a modelled bladder contraction (part A) and a plot of  
381  $(dP/dt).P^{-1}$  as a function of  $P$  (Part B). The plot can be fitted by a hyperbolic function  
382 equivalent to a Hill force-velocity relationship;  $(p+a)(v+b)=b(p_0+a)$ . Here  $a$  and  $b$  are  
383 constants,  $P_0$  is the maximum value when  $v_{CE}$  (equivalent to  $(dP/dt).P^{-1}$ ) is zero and the  
384 intercept on the  $(dP/dt).P^{-1}$  axis is equivalent to maximum  $v_{CE}$ .

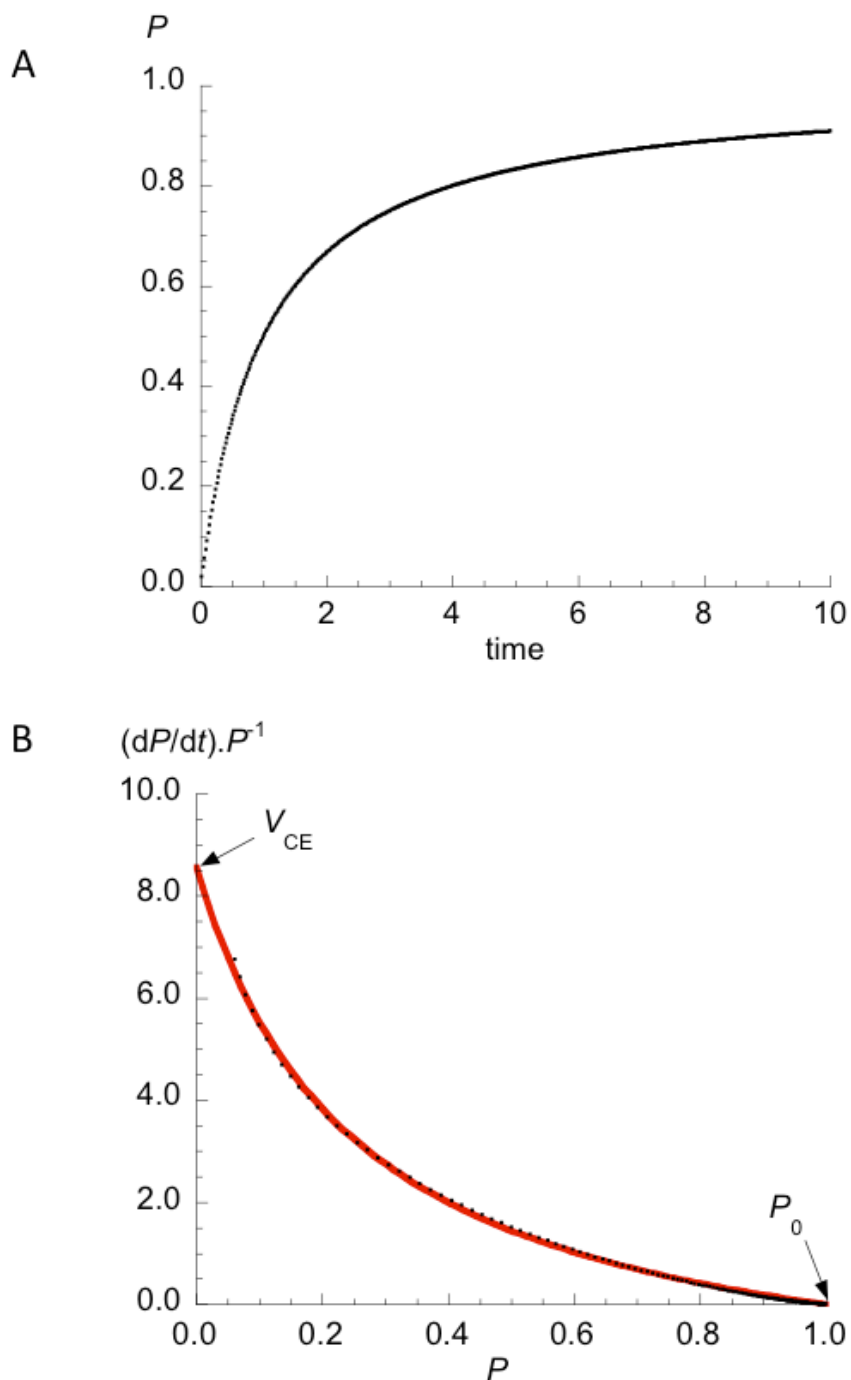
385

386 1. Urschell et al. 1970. Circulation 42 (suppl III): 111-115, 1970.

387

388

389



391 Supplementary Figure 1. Derivation of the method to estimate muscle contractility. A model  
 392 contraction (part A) and a derived  $(dP/dt).P^{-1}$  vs  $P$  (part B). The line in part B is fitted to the  
 393 equation of the hyperbola  $(p+a)(v+b) = b(p_0+a)$  and extrapolated to the  $P$ -axis and  $(dP/dt).P$   
 394  $^{-1}$  axis to yield respectively values of  $P_0$  and  $v_{CE}$ .



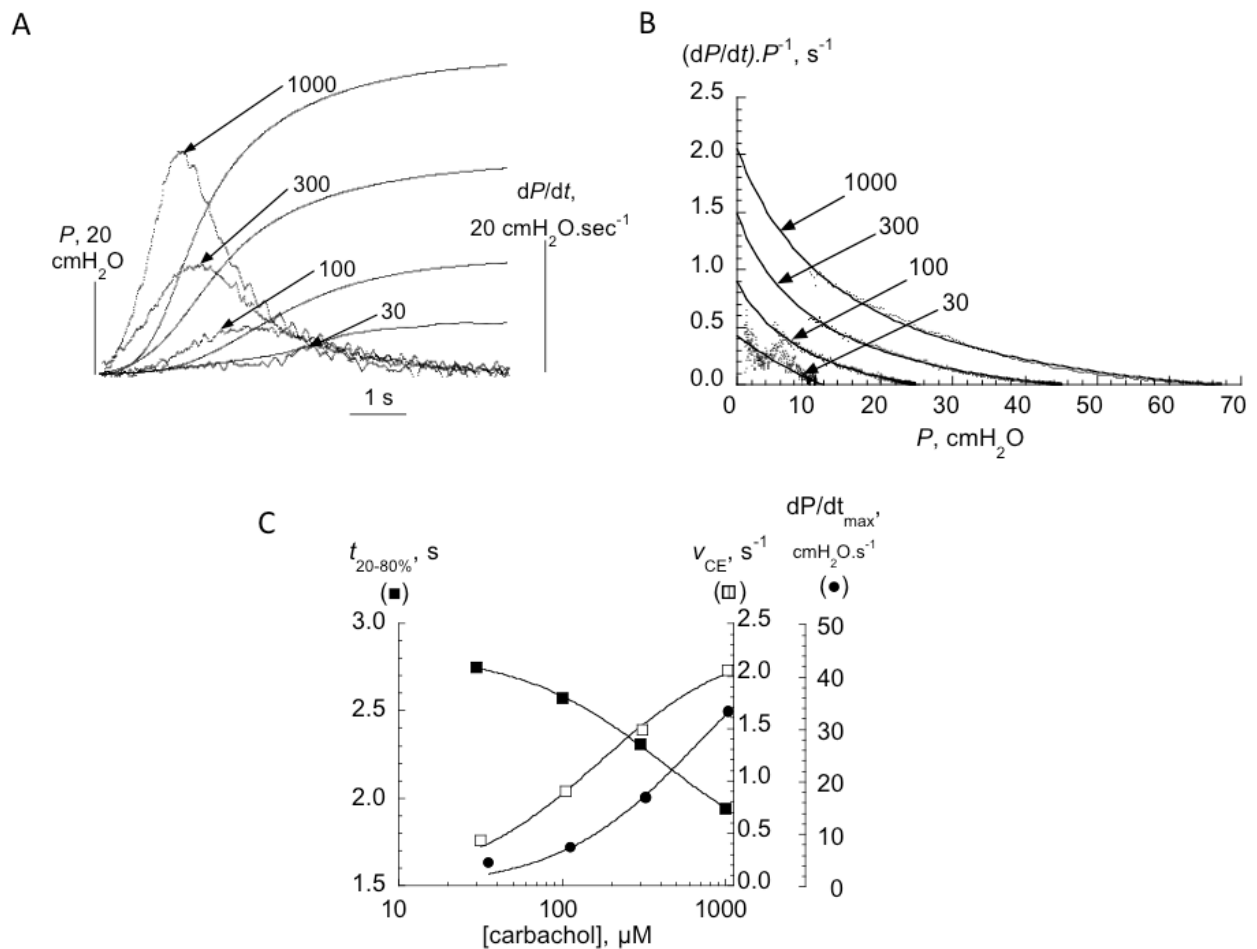
396 Validation of the analysis using an inotropic agent with isolated perfused pig bladders.  
397 Experiments used arterially perfused pig bladders to demonstrate the action of a known  
398 inotropic agent carbachol, added to the perfusate, on intravesical pressure and also on  $v_{CE}$  and  
399  $P_0$ . The bladder was held at constant volume to generate isovolumetric contractions, as  
400 measured in the human subjects. Pressure recordings were less prone to external  
401 interference than recordings with humans and so provided a convenient validation model.

402

403 *Methods.* Female pig bladders were obtained at a local abattoir. The distal abdominal aorta  
404 and branches supplying the bladder were carefully dissected free. The bladder and associated  
405 vasculature were then excised and perfused with ice-cold Ca-free Krebs' solution at 4°C to  
406 drain blood from the organ and transported to the laboratory in cold (4°C) solution for  
407 immediate use. *Ex vivo* intravesical pressure recordings were made from arterially-perfused  
408 pig bladders at 37°C and filled to 150 ml.<sup>1</sup> Isovolumetric contractions were elicited by  
409 injections of carbachol-Krebs's into the perfusion-line. Digitised recordings were recorded at  
410 10 kHz and retrieved at 0.4 kHz for analysis. The composition of Krebs's solution was (mM):  
411 NaCl, 118.3; NaHCO<sub>3</sub>, 24.9; KCl, 4.7; MgSO<sub>4</sub>, 1.15; KH<sub>2</sub>PO<sub>4</sub>, 1.15; CaCl<sub>2</sub> 1.9; D-glucose, 11.7,  
412 gassed with 95% O<sub>2</sub>/5% CO<sub>2</sub>, pH 7.38 ± 0.01, 36±1°C).

413

414 *Results.* Pressure transients were recorded, at a constant intravesical volume of 150 ml, in  
415 response to the contractile agonist, carbachol (30-1000 µM). Supplementary Figure 2A shows  
416 superimposed rising phases of pressure transients and their first derivatives in response to  
417 carbachol. Corresponding  $(dP/dt).P^{-1}$  vs  $P$  plots are in part B and show that increased  
418 contractions and  $dp/dt_{max}$  values with rising carbachol concentrations were mirrored by  
419 increases of  $v_{CE}$  and  $p_0$  (part B). Part C shows the concentration-dependence of  $v_{CE}$ , and  
420  $dp/dt_{max}$ . In addition,  $v_{CE}$  was inversely related to values of  $t_{20-80}$ , as in human data.



Supplementary Figure 2. Contractility variables in the *ex vivo* pig bladder. A: Isovolumetric pressure traces from perfused pig bladder during bolus injections of carbachol (30-1000  $\mu\text{M}$ ): also shown are the first derivative ( $dp/dt$ ) of the traces. B: plots of  $(dP/dt) \cdot P^{-1}$  as a function of  $P$  for the traces in part A. C: carbachol dose-dependence of  $v_{\text{CE}}$  (open squares) and  $dP/dt_{\text{max}}$  (closed circles) and  $t_{20-80\%}$  (closed squares). Lines are those of best-fit.

## Reference

- Parsons BA, Drake MJ, Gammie A, Fry CH, Vahabi B. Validation of a functional, isolated pig bladder model for physiological experimentation. *Front Pharmacol.* 2012 Mar 30; 3: 52.